in weight. Fourteen days following the first stage operation he was given ether and the prostate enucleated. Within twenty minutes the patient had been returned to bed and was in excellent condition. On the fifth day following the operation the temperature and respiration were normal and the pulse was 110 and somewhat irregular. There was no hemorrhage; the urine was not bloody. The large suprapubic drainage tube had been removed and a smaller one had been instituted. The patient was rational. Respirations were thirty. There was a certain amount of dyspnea present. Twelve hours later the dyspnea was very great. The pulse was very rapid and thready. The apex-beat was diffuse and the patient had labored breathing. There was edema of the ankles present. Patient was given stimulation and water was increased as much as he would take. The dyspnea increased rapidly and the heart action became weaker and death occurred the following day. At the autopsy the diagnosis was mitral endocarditis with insufficiency. Chronic myocarditis, edema of both lungs, chronic passive congestion of the liver and spleen and chronic nephritis.

The case records which are quoted here are typical of their class and the autopsy findings are representative of what is found as a cause of death. No effort has been made to determine the mortality rate. The death-rate from prostatectomy has not been higher than with other individuals and institutions. More recently there have been fewer deaths, and I have attributed it to a better understanding of the presence of infection in the form of pyclonephritis which threatens prostatics, and to a prolonged, free, suprapuble drainage, and treatment for infection in their preparation for operation.

URINARY ANTISEPSIS: A STUDY OF THE ANTISEPTIC PROPER-TIES AND THE RENAL EXCRETION OF 204 ANILIN DYES,1

BY EDWIN G. DAVIS, M.D., OMAHA, NEB.

In 1915 Hinman² pointed out the inefficiency of the several drugs at present in common use as urinary antiseptics and concluded that there is no known drug ideally suited for this purpose or even approaching the ideal. These observations are borne out by the experience of everyone who has had to deal with chronic infections of the urinary tract and also by the absence in the literature of conclusive experimental or clinical evidence of the fitness of any drug

² Urinary Antisepsis: A Clinical and Bacteriological Study, Jour. Am. Med. Assn., 1915, lxv, 1769.

¹ Investigations were carried on in the laboratory of the University of Nebraska College of Medicine, with the aid of an appropriation made by the United States Interdepartmental Social Hygiene Board.

for this purpose. The ideal internal urinary antiseptic should be chemically stable and relatively non-toxic and non-irritating; should be antiseptic in high dilution in urine as well as on agar (regardless of the reaction of the former), and should be eliminated in high percentage by the kidney without injury to the body. Clinically there is no such drug known.

For the purpose of urinary antisepsis, urotropin is the most widely used, and likewise the best suited available drug, on account of its well-known and proved action of liberating formalin in the urine. Urotropin, however, has very definite limitations, owing largely to the necessity for an acid urine for the liberation of formalin, and this becomes an insurmountable obstacle in those urines infected with alkalinizing organisms, such as the micrococcus urea or bacilli of the proteus group. As is well known, by the administration of acid sodium phosphate it is possible to cause a temporary slight increase in urinary acidity.

Henderson and Palmer' have shown, however, that even after the administration of 10 grams of acid sodium phosphate at a single dose there follows only a slight increase in the hydrogen ion concentration, and in no case were they able to produce a urinary acidity greater than that which they had commonly observed in patients to whom no drug had been administered. On the other hand to produce and maintain a relatively large variation toward the alkaline end of the hydrogen ion scale, by the use of sodium bicarbonate, is comparatively simple. Therefore, granting that the efficiency of a given urinary antiseptic must necessarily be dependent upon the reaction of the urine, a drug efficient in alkaline urine only would be of greater practical value than is urotropin.

Another recognized limitation to urotropin is the time necessary for formalin liberation (Burnam'), which destroys the value of the drug at the kidney level and also in the bladder in eases in which for any reason there is rapid emptying or where a fistula exists. The main value of urotropin lies in its use as a prophylactic before instrumentation.

Summary of Previous Studies. Previous publications record the results of investigations with reference to the synthesis of an internal urinary antiseptic, carried on at the Brady Urological Institute, Johns Hopkins Hospital, in cooperation with Dr. Edwin C. White, ehemist for that institution. (Davis, Davis and White, Davis,

³ On the Extremes of Variation of the Concentration of Ionized Hydrogen in Human Urine, Jeur. Biol. Chem., 1913, xiv, 81.

⁴ An Experimental Investigation of the Value of Hexamethylenamin and Allied Compounds, Arch. Int. Med., 1912, x, 324.

Urinary Antisepsis: A Study of the Antiseptic Properties and Renal Excretion of Compounds Related to Phenoleulphonephthalein: Preliminary Report, Jour. Am. Med. Assn., 1918, 1xx, 581.

Urinary Antisepsis: Further Studies of the Antiseptic Properties and Renal Exerction of Compounds Related to Phenolsulphonephthaloin, Jeur. Urel., 1918, ii, 107.

White and Rosen?). Here an attempt was made to correlate chemical structure with the antiscptic properties and the renal exerction of the compounds studied, most of which were synthesized for this special purpose, with the hope that the introduction of certain groups into the molecule would produce certain desired properties. The study was limited largely to compounds related to phenolsulphone-phthalein, because of the well-known extraordinary "renal affinity" possessed by this compound and because of its non-toxicity. Some interesting results were obtained which may be briefly summarized as follows:

1. It was possible to establish a certain relationship between chemical structure and renal exerction and to predict the exerction of molecules of certain structure, particularly those of the xanthone group. The halogenation of these compounds interfered with exerction.

2. Many of these compounds, non-toxic, exercted in the urine and antiseptic in water, lost this latter property when tested in voided urine.

3. One compound, chlor-mereury fluorescein, experimentally possessed all of the required properties, and when administered intravenously in minute dosage (5 mgm.) to dogs and rabbits caused the secretion of antiseptic urine for a definite period of time without

injury to the animal.

Clinical investigation of this drug has not been carried out on account of its mercury content, although it was shown that in dogs the single lethal dose was forty times that necessary to cause the secretion of antiseptic urine. Chlor-mercury fluoresecin, therefore, approaches the ideal in that (a) it is antiseptic in high dilution in either acid or alkaline urine; (b) it is exercted by the kidney with a rapidity as great as is phenolsulphonephthalein, and (c) experimentally efficient dosage may be administered without toxicity. Chlor-mercury fluorescein is an organomercury phthalein derivative in which the mercury is present in non-ionic form.

4. Continued experiments along the same lines (Davis and White⁸) have shown that aeriflavin and proflavin are antiseptic in high dilution in urine (particularly in alkalin urine) and that intravenous administration of minute dosage (5 mgm. per kilo) to rabbits causes secretion of urine, which is antiseptic for a definite period of time without injury to the animal. Rabbit urine is normally usually alkaline. Failure to produce antiseptic urine in dogs with corresponding dosage of the same drug was probably due to the fact that

dog urine is usually acid (average about ph, 6).

Preliminary Report, Jour. Urol., 1918, ii, 299.

⁷ Urinary Antisepsis: The Secretion of Antiseptic Urine Following the Intravenous Administration of an Organomercury Phthalein Derivative, Jour. Urol., 1918, ii, 277.
⁸ Davis, E. G., and White, E. C.: Urinary Antisepsis: The Secretion of Antiseptic Urine Following the Intravenous Administration of Acriflavin and Proflavin.

Possibilities Offered by Anilin Dyes. The following record summarizes the results of an investigation of the antiseptic properties and the renal excretion of 204 anilin dyes, the scope of the work being limited and guided, not by the ehemical structure of these compounds but only by the number available. This investigation was carried on in the laboratories of the University of Nebraska College of Medicine with the aid of an appropriation made by the

United States Interdepartmental Social Hygiene Board.

The anilin dyes were chosen for study (1) because of the large number of these compounds available, (2) because of their color and bence their ready detection and quantitative estimation in the urine and (3) because, through the work of many observers (notably, Churchman, Krumwiede and Pratt, 10 Simon and Wood, 11 Kligler, 12 Graham-Smith¹³), the antiseptic properties of certain anilin dyes have become well known and therapeutic possibilities in this field have been indicated. Furthermore a consideration of the history of the development of the various tests of renal function (Thomas and Birdsal14) will eall to mind that there are several dyes (fuelsin, rosanilin, indigo-carmin, uranin, trypan blue and others) which have been used to measure the functional activity of the kidneys, and which are therefore known to be exercted without injury to the patient. The staining and penetrating properties possessed by many anilin dyes likewise suggest suitability of this type of compound for medication of the urethral mucosa. This investigation was not undertaken without due realization of the handicap presented by impurities in commercial samples of anilin dyes.

Method of Investigation. Considering the large number of dyes to be studied it was advisable to select a few by preliminary test on agar, thus ruling out many as being unworthy of further investigation. The remaining few were then studied in regard to their antiseptic value in urine, their toxicity, their renal exerction and in regard to their ability to cause the secretion of antiseptic urine following intravenous administration. Finally, those few which were found to be particularly efficient against the stapbylococci were tested on special media against the gonoeceeus. (Tables showing results with the gonococcus will appear in a subsequent publication.) The investigation was therefore divided into five stages as follows:

⁹ The Specific Antiseptic Action of Gentiao Violet Corresponding to Gram, Jour. Exper. Med., 1912, xvi, 221, 822.

¹⁰ Observations on the Growth of Bacteria on Media Containing Various Aniliu Dyes, Jour. Exp. Med., 1914, xix, 20 and 501.

¹¹ The Inhibitory Action of Certain Anilin Dyes upon Bacterial Development, AM. Joun. Med. Sc., 1914, exlvii, 247.

¹² A Study of the Antisoptic Properties of Cortaio Organic Compounds, Jour. Exp. Med., 1918, xxvii, 463.

¹³ Some Factors Influencing the Actions of Dyes and Allied Compounds on Bac-

toria, Jour. Hyg., 1919, xviii, i. ¹⁸ Comparativo Results of Various Functional Renal Tests, Based on a Series of Cases, Jour. Am. Med. Assn., 1917, lxix, 1747.

1. Antiseptic Values on Agar. Determination of the antiseptic strength of the entire list of dyes on agar against B. eoli, Staphylococcus albus and Staphylococcus aureus.

 Antiseptic Values in Urine. Determination of antiseptie strength of selected dyes in both acid and alkaline urine against B.

coli, S. albus and S. aureus.

3. Taxicity and Excretian. Determination of toxicity and renal exerction in rabbits of dyes shown to have antiscrtic value in voided urine.

4. Experimental Urinary Antisepsis. Determination of the antiscptic value of the urine of rabbits which had received intravenous injections of dyes previously shown to be non-toxic and excreted.

5. Inhibitian of Gonacoccus. Determination on special media of the antiscptic strength against the gonococcus of those dyes which had been shown to inhibit the staphylococcus in high dilution.

1. Antiseptic Values on Agar. Preliminary antiseptie tests were carried out on the entire list of dyes, using agar neutral to phenolphthalein and of the following composition:

Agar													15 000
Pantona (Witte)		•	•	•	•	•	•	•	•	•	•	•	to Rm.
Peptone (Witte)	•	-	•	•	•	•	•		•		•		10 gm.
MICH DATING (LIGHE)	-												E
continu cutoride				-									E ~~~
Water							-	-	-	•	•	•	1000
	•	•	•	•		•	•	•	•		•		TOOR G'G"

Since the colon bacillus is by far the most frequent invader of the urinary tract, this organism was chosen together with S. albus and S. aureus. The agar was autoclaved in test-tubes in 9 c.c. amounts, after which 1 c.c. of an aqueous solution of the dye was added, the latter solution being at a concentration ten times that desired for the final dilution. Each dilution was then plated, cooled and inoculated by three parallel strokes from twenty-four-hour broth cultures of the three above-named organisms. No concentrations greater than 1 to 1000 were used, all dyes not showing antiseptic properties at this concentration being discarded. The selective antiseptic action against various organisms which Churchman has described, with particular reference to gentian violet, was exhibited by no less than 44 dyes, and in every case it was the colon bacillus that survived, while one or the other of the staphylocoeci (usually both) failed to grow. (See Figs. 1, 2 and 3.)

2. Antiseptic Values in Urine. As previous publications on this subject have indicated the possession of antiseptie properties by a drug when diluted in water or in agar is no indication whatever of its antiseptie value when diluted in urine. Many of the sulphone-phthaleins which were germicidal in high dilution in water lost this property when diluted in urine in a test-tube and even permitted the growth of organisms in urine when in relatively high concentration. In determining the antiseptie value in urine of the dyes selected by preliminary test on agar it was therefore desirable to

make the various dilutions with voided urine, since any drug for the above purpose would be useless unless effective in this medium. Furthermore, it was necessary to try out each dye in both acid and alkaline urines, since the ideal drug should be efficient regardless of the urinary reaction.

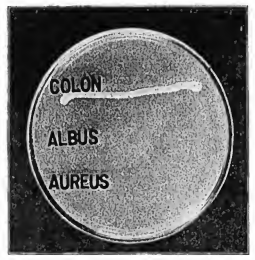


Fig. 1.—Photograph of agar plate containing chrysoidin-Y (1 to 1000), showing selective action of the dye in permitting growth of Bacillus coli and inhibiting Stuphylococcus ulbus and Staphylococcus autors.

In order to have available each day urine of definite acid and alkaline reaction it was necessary to titrate specimens of voided urine with tenth normal sodium hydroxide and tenth normal hydrochloric acid until definite degrees of hydrogen ion concentration were reached, as determined by the colorimetric method—that is, by comparison with a standard hydrogen ion scale made up with solutions of buffer salts colored by the sulphonephthalein series of indicators. (See publications of Clark and Lubs¹⁵ and Shohl and Janney. (So On the acid side of the scale it was arbitrarily decided to use urine titrated to pb, 6.4, which Henderson and Palmer have shown to be slightly less acid than the average reaction of normal urine. In

n Colorimetric Determination of Hydrogen Ion Concentration, Jour. Bacteriol., 1917, ii, 1.

³⁶ Growth of Bacillus Coli in Urine at Varying Hydrogen Ion Concentrations, Jour. Urol., 1017, i, 211.

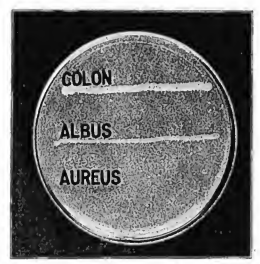


Fig. 2.—Photograph of ngar plate containing phloxin-P (1 to 1000), showing selective action of the dye in permitting growth of Bacillus coli and Staphylococcus albus and inhibiting Staphylococcus areus.

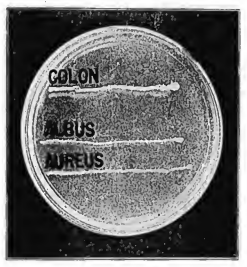


Fig. 3.—Photograph of control plate of drug-free agar, showing profuse growth of all three organisms.

order to obtain alkaline urine a sample of the same specimen was titrated to p_b, 7.6, an end-point arbitrarily chosen so that the reaction of the specimens of urine used from day to day would not vary.

Dilutions of the dyes were made in sterile test-tubes, using acid urine for one series of dilutions and alkaline urine for another. Each dilution was inoculated with one loop of a twenty-four-hour

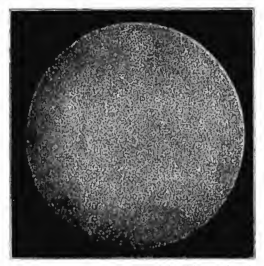


Fig. 4.—Photomicrograph (low power) of typical field in agar plate, showing absence of colonies and proving that the corresponding urine tube had contained a concentration of dye sufficient to kill the organisms during the twenty-four-hour incubation period.

broth culture of Bacillus coli in one series of experiments and with the Staphylococcus albus in another. After an incubation period of twenty-four hours (sufficient time to permit either growth or death of the organism), 0.1 c.c. was transferred from each tube to melted agar and plated. Those plates remaining sterile after incubation (Fig. 4) proved that the urine in the corresponding tubes had contained a concentration of the dye sufficient to kill the organisms within twenty-four hours. Those plates showing a few scattered colonies after incubation (Fig. 5) proved that the concentration of the dye in the urine had been sufficient to cause an arrest in the development of the organisms. This is the "bacteriostatic" action of antiseptics referred to by Hinman and should be

sufficient to control urinary infection provided the administration of the drug is continued. Finally, plates in which countless numbers of colonies developed (designated in the tables by the infinity sign ∞) proved that the concentration of the drug had been insufficient to prevent growth of the organism (Fig. 6).

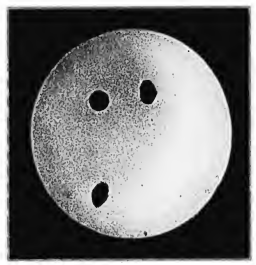


Fig. 5.—Photomicrograph (low power) of typical field in agar plate, showing a few scattered colonies and proving that the corresponding urine tube had contained an inhibitory or bacteriostatic concentration of the dye.

A consideration of Table II permits of several general conclusions. In keeping with the well-recognized clinical frequency of colon bacillus urinary infections, and with the stubbornness of such infections, is the hardiness which this organism displays in urine in vitro in spite of the presence of antiseptic dyes. Out of a total of 204 dyes studied only 24 prevented the growth of the colon bacillus in urine in a dilution of 1 to 1000. Of these several were effective in alkaline urine only. A 1 to 1000 solution is a relatively high concentration, considering the extreme dilutions (several greater than 1 to 1,000,000) at which these same dyes are effective against staphylococci in the same media at the same reaction Table II further shows that it is the general rule that these dyes are antiseptic in higher dilution in alkaline than acid urine. This fact might well prove to be of clinical importance, since the artificial production and maintenance of an alkaline urine is a relatively simple matter.

TABLE I.—RESULTS OF PRELIMINARY TESTS ON AGAR OF ANTI-SEPTIC VALUE OF THE ENTIRE LIST OF DYES AGAINST COLON BACILLUS (C), STAPHYLOCOCCUS ALBUS (AI) AND STAPHYLO-COCCUS AUREUS (Au). ALL DILUTIONS ARE I TO 1900. O =GROWTH. — = NO GROWTH.

Name.	No.	Az	ıtise reng	ptic	Name.	No.		ntise	
***************************************		C.	Al.	Au		1.0.	c.	AI.	Au.
Nitrose. Nnphtholgreen B	4*	0	0	0					
Nitne Mnrtius yellow .	٠ 6	-	-	-	Nitno. Noplithol yellow S	7	0	0	0
STILBEN. Goldeo yellow	9	0	0	0					
Pyrazelen. Flavazin L Flavazin S	19 20	8	8	8	Pyrazolon. Tartrazin	23	0	0	0
Aze (Mencaze). Chrysoidin R Soudan I Poncenu 4 GB Orange G Chrometropo 2R Orange III Orasellior Mixt. Chrysoidin Mixt. Chrysoidin Srilliant orange O Brilliant orange O Brilliant orange R Brilliant orange R Palatine scarlet A Ponceau R Ponceau 3R Ponceau 3R Ponceau 3R Ponceau 3R Azocosin G Azocosin Fast red	33 34 36 37 38 40 47 53 69 70 70 81 82 83a 83b 94b 112	0000000000000000000	11000000000000000000	1100000000000000000	AZO (MONOAZO). Erika B oxtra Victoria yel. O Retanii yel. extra Fast yellow Brilliont yel. S. Chrysoin Orango I Azofuelsin Fast yellow Orango T Nnphthyl nuino br. Azo rubiii Fast red Fast red Fast red Fast red D Coelenille red A Fhat brown 3B	121 122 134a 134b 137 142 143 144 145 140 151 163 160 168 169 172	000000000000000000	. 000100000000000000	000100100000000000
Azo (Diazo). Resorcia brown Fast brown G Fast brown Bluo black Folotine black Folotine black Brilliont eroceiu Soudan IV Fast scarlet Crocein scarlet Crocein scarlet Crocein start Bismark brown	211 212 213 215 229 227 232 248 249 255 283	00000000000	00000000001	10000000001	Azo (Diazo) Vesuvin B Azonlazarin Bord. W. Brilliant yellow Benzopurpurin 4B Benzopurpurin 6B Ressaurini Azo blue Dianil blue B Trypaublue Cirysanin R Bovorion blue	284 291 393 363 364 372 377 380 391 394 450	0000000000	01000000001	01000000001
AURAMINE. Auromino	493	-	-	-	AURAMINE Auramine G	404	o		_
TRIPHENYLMETHANE. Malnchite green Brilliant green Lt. green SF bluish Erioglaucine A Fluchsin Red violet 5R ex. Methyl violet Bcrystal violet Methyl violet Methyl violet Methyl violet Methyl violet Methyl violet Methyl violet Aniliae blue	495 499 594 505 512 514 515 516 517 518 519	110000111100	110001111100	110001111100	Thi-HENYLMETHANE. Victoria blue 4R Fuchsin S Red violet 5RS Acid violet 4BN Acid violet 54B Methyl blue Soluble blue Amalga green B Patent blue Patent bluo Cyanol extrn	522 524 525 527 527 538 538 542 543 546	000000000000	100100100000	100100100000

^{*} Numbers correspond to Gustav Schultz, Farbstofftabellen, 1914 (5th edition).

TABLE I .- Centinued.

Name.	No.	An	tiser	ntic th.	Name.	No.	An	tiser	otic th.
- Tunio		C.	Al.	Au.			c.	At.	Au.
DIPHENYLNAPHTHYL- METHANE, Victorio blue R Victoria blue	558 550	00	0 -	0	Diphenylnaphthyl- methane, Wool green S	566	0	0	_
XANTIONE. Redamine S Rodamino 0G extra Rodomine G extra Rodomine B Rodamino 3G Sulphoredamino B Fast ocid violet B Fast ocid violet Ag Reddrosamino Fast ncid bluo Chryselin	576 571 572 573 576 570 580 582 583 584 586	00000000000	11111000000	11111000001	XANTHONE. Eosin Methyleesin Eosin BN Erythrosin G Erythrosin Phloxine P Phloxine P Phloxine P Phloxine R Ross bengale Rose bengale	587 588 590 501 502 593n 593b 596 597a 507b	0000000000	0100010111	0101111111
ACRIBINE. Benzeflavine Phesphine.	665 606a	0	_	=	ACRIDINE. Phosphine N Rheonin	606b 667	00	=	=
Quineline. Quineline yellew	612	o.	0	0					
Thie Genzenyl. Thieflavine S Thieflavine S	615a 615b	00	00	00	Primuline	616	0	0	0
OXAZINE, Meldela's blue New blue B New methylene blue	649 656 651		- - -	=	Oxazine. Methylene blue	659	-	-	-
THIAZINE. Thie carmine	662	0	0	0					
Azine. Flavindulin O Neutral red Indulin scarlet Safranin T Safmuin OW	668 670 671 679 683a	10000	1011	1011	AZINE. Safranin MN Nigrosine Indulin NN Indulin	683b 608 699 766	0000	1000	1000
MISCILLANEOUS, Searlet of R Diphenyloimine eminge Crystal scarlet Biebrich sanlet Creecine 3BX Guinea green Quinone Pentneyl sub. blue 5RX Pontacyl ponceau Pon. sul. acid blue R Pontneyl nzo flavine Pen. sul. black 2B Pon. bline black SX Pentamine violet X Pontamine red B Pontamine blue AX Pontamine sky blue 5B Pentamine blue AX Pontamine blue AX Pontamine blue B Ponta, bline blue SR Pentamine blue B Ponta, dieze bl. BH Pontachrone black F Pontachrone bl. 6BX Ilydron blue G Stulphegoeo Bordeau B Gentian violet and service of the ponta of the service black F Pontachrone bl. 6BX Ilydron blue G Stulphegoeo Bordeau B Gentian violet		000000100000000000000000000000000000000	00000010000000000000001001	00000010000100000000001001	Miscellaneous. Pontacyl pr. Congo red Searlet B Pontamine brewn R Pontamine green GX Pontamine green GX Pontamine black EX Pontamine black EX Pontachrone yellew Pontamine blue ZX Safranine T extra Pontamine yel, SX Primuline Pontamine yel, SX Primuline Pontamine of star Pontamine of star Pontamine of star Richibert of star Pontamine of star Richibert of star Sulphegene ind. blue G Aurine Sulphegene or, L Titionel black XX Sulphegene any blue Hydron blue R Sulphegene br. G		0 00000001000000000000000	0 000000011000000010000000	0 000001011000000101100000
STANDARD ANTISEPTICS. Mercuric chlorido Phenol	::	- 0	- 0	- 0	STANDAND ANTISEPTICS. Silver nitrate Ethyl alcohol		0	0	0

TABLE II.—DETERMINATION OF ANTISEPTIC VALUE IN BOTH ACID AND ALKALINE URINE OF 28 DYES SELECTED BY PRELIMINARY TEST ON AGAR FROM THE ORIGINAL 204, AS SHOWN IN TABLE I.

					W	ntiseptie stre	Antiseptie strength in urine.			
				Colon !	Colon bacillus.			Staphyloce	Staphylococcus albus.	
Group.	Dye.	No.	Aeid (pr.	Aeid urino · (pn, 6.4).	Alkalin (ps.	Alkaline urine (pri, 7.6).	Acid (ps,	Acid urino (pg. 6,4).	Alkalin (pu,	Alkaline urine (pu, 7.6).
			Inhihits devel. I to	Permits growth. I to	Inhihits devel. I to	Permits growth,	Inhihits devel. 1 to	Permits growth.	Inhihits devel,	Permits growth.
Nitro Azo Auramine Triphenyl-	Martius yellow Chrysoidin Y Chrysoidin R Victoria yellow Auramine Auramine G	33 34 34 134b 493 494	1,000	10,000 10,000 1,000 1,000 1,000	1,000	10,000 1,000 1,000 3,000 5,000	10,000* 10,000* 30,000 1,000 1,000		10,000* 50,000* 1,000 1,000	10,000
ethane .	Mal green Brill, green Ruchsin Rod viol, 5R ox. Methyl viol, B	55129 51129 51154 5154	5,000* 1,000* 3,000	1,000	1,000* 1,000* 1,000* 1,000	10,000*	900,000 100,000 1,00,000	1,000,000 1,000,000 300,000	1,000,000* 900,000 100,000 700,000* 1,000,000*	1,000,000
Xanthone	Methyl viol. Ethyl viol. Victoria blue	512 522 571 571 571	1,000*	000;1	8,000	5,000 1,000 1,000	3,000,000 3,000,000 10,000 30,000	rD.	000,000 000,000 000,000 000,000	ō'-ţ
Acridin	Roso beng. cx. Benzoflavine Rheonin	597b 005 007	1,000*	000,1	1,000*		30,000 30,000 30,000		100,000	
Azine	Flavindulin Indulin scar Safranin T Safranin OW	008 671 679 683a	1,000	1,000	1,000	00000	900,000 1,000 10,000	Ħ.	200,000 200,000 200,000 200,000	
Miscellaneous .	Safrann MN Acriffayine Safranin T ex. Gentian violot	083b	5,000	1,000	1,000* 100,000 2,000*	200,000	10,000 30,000 100,000	100,000 100,000 20,000	100,000	200,000 100,000 900,000
tics .	Phenol Hg. bichlorido Ag. nitrato	:::	10,000	30,000 30,000	10,000	30,000	10,000	30,000	10,000	

3. Toxicity and Excretion. Twenty-seven dyes were selected from Table II as seeming worthy of further study as to toxicity and excretion. Rabbits were given intravenous injections of from 5 to 25 mgm, per kilo and the urine collected at intervals and examined for the dve. Only in those cases in which the excretion was strikingly rapid and complete was an attempt at quantitative colorimetric estimation made. As shown in Table III, several dyes (malachite green, brilliant green, crystal violet, ethyl violet, victoria blue and others) were exceedingly toxic, causing convulsions and death within a few minutes. Autopsies showed a varying and bizarre sclective distribution of the different dyes through the various tissues of the body. Several dyes (for instance chrysoidin R, crystal violet, rhodamin 3G, benzoflavin, indulin scarlet and others), though not fatal, were ruled out on account of hematuria or hemoglobinuria following minute dosage. Particular attention is called to the triphenylmethane group, of which many showed antiseptic properties in extreme dilution in urinc. Dyes of this group, however, were also the most toxic, and those few which did not injure the rabbit failed to appear in the urine. (See Table III.)

TABLE III.—EXCRETION AND TOXICITY. RESULTS OF INTRAVENOUS INJECTIONS IN RABBITS OF DYES SHOWN IN TABLE II TO POSSESS ANTISEPTIC VALUE IN URINE.

			Effec	os asimal.	Renal excretion.			
Group.	Dye.	No.	Dose, mgm. per K.	Result.	Dose, mgm. per K.	Result.		
Nitro , , .	Mnrtius yellow	6	10	None	10	Me dernte.		
Azo	Chrysoidin Y	33	20	None	20	Mnrked.		
	Chryseidin R	34	20	Hematuria	20	Moderete.		
	Victorin yellow	134b	20	None	20	Moderate.		
Aurnmine .	Auromino	493	10	None	10	Medernte.		
	Auramine G	494	10	None	10	Nene.		
Triphenyl-	•			2.020		21020		
methane .	Malachito green	495	20	Lethel	4	None.		
	Brilliant green	499	10	Lethnl	4	None.		
	Fuchsin	512	20	None	20	None.		
	Rod viol. 5R oxtre	514	20	Lethel	15	None.		
	Methyl violet B	515	30	Apuria	20	None.		
	Crystal violet	-516	4	Hematuria*	4	Nono.		
	Methyl violet	517	40	Lethnl	20	None.		
	Ethyl violet	518	20	Lethal	8	None.		
	Victoria blue	522	8	Letbnl	8	None.		
Xanthono .	Rhod, 0G extra	571	20	Lethel	10	Moderate.		
	Rhod. 3G	576	20	Hematuria	liŏ	Moderate.		
	Rose beng, extra	597b	20	Nono	20	None.		
Acridin	Benzoflavine	005	10	Homaturia	4	Slight.		
	Rheonin	607	20	Nono	20	None.		
Azino	Flavindulin	068	40	Lethal	20	Doubtful.		
	Indulin scar.	671	10	Hematuria	10	Marked.		
	Safranin T	679	20	None	20	Moderate.		
	Safrnnin OW	683a	20	None	20	Marked.		
	Safrnnin MN	683b	20	None	20	Marked.		
Miecellaneoue	Safranin T extra		30	None	20	Marked.		
	Acriflavine		20	None	5	Mnrked.		
	Proflavine .	:::	20	None	5	Mnrked.		

^{*} Hemnturia lasted three days.

4. Experimental Urinary Antisepsis. Table III shows that of the total of 204 dyes studied there remained only 13 which were antiseptic in urine (in vitro), which were excreted by the kidney after intravenous administration and which exhibited no toxic properties following moderate dosage (about 20 mgm. per kilo). These dyes are listed in Table IV. It then remained to attempt to demonstrate antiseptic properties in the nrine of rabbits following the intravenous administration of these drugs; that is, to determine whether passage through the blood stream and kidney would interfere with the antiseptic properties, and whether sufficient dosage could be safely administered to cause adequate concentration in the urine.

TABLE IV.—EXPERIMENTAL URINARY ANTISEPSIS. RESULTS OF ATTEAUPTS TO CAUSE THE SECRETIONOF ANTISEPTIC URINE BY THE INTRAVENOUS ADMINISTRATION TO RABBITS OF DYES, SHOWN IN TABLES II AND HI TO BE ANTISEPTIC, EXCRETED AND .RELATIVELY NON-TOXIC. C = COLON BACILLUS. AI = STAPHYLOCOCCUS ALBUS. \(\omega = AN INFINITE NUMBER OF COLONIES. \)

				ur	ino w	pia hich	had r	ntain	ing	0.1 c	ed in .e. of lnocu- hours.
Group.	Dye.	No.	Dose, ingm. per K.	obti ji be	rino nined ust foro etion.	obt	rine ained oura ter etion.	obti 6 h	Urine obtained 0 houra alter injection,		rine nined ioura ter etion,
•				C.	Al.	C.	Al.	C.	Al.	C.	Al.
Nitro	Martius yellow	0	10	8	80			- 8	<u> </u>		80
Azo	Chrysoidin Y	33	20	80	8		6		80	, w	60
	Chrysoidin R	34	20	80	6	8	80	· ·	· ·		8
	Victoria yellow	134b	20	8	8	89	m	00		~ ~	
Auramino .	Auramino	403	10	80	80	60	6	80	80	8	60
Xanthone	Rhod. 6G extra	571	10	80	80	ထ			80		80
	Rhod. 3G .	576	10	œ	ω	œ	80	80	00	60	
Azine	Flavindulin	668	20	8	80	œ		80	8	80	00
	Indulin scar	07 I	4	80	8	œ		i	80	60	80
	Safranin T	670	20	80	έω	80	80	œ .	80	80	BD
	Safranin OW	083a	20	œ	ω .	œ	80	80	00	60	60
	Safranin MN	083b	20	. ۳۵	80	00	8	80	80	60	60
Miscellaneous	Safranin T extra		30	œ	60	ထ	60	60	80	60	œ
	Aeriflavine		10	œ	00	0	0	0	0.1		80
	Proflavine	<u> </u>	10	80	ω	0	0	ŏ	ő	80	80

In general the method of procedure was to compare the antiseptic properties of several specimens of urine, obtained by eatheterization from a given rabbit before and at intervals of from one to several hours after drug administration. The necessity for a control urine has been pointed out in a previous publication (Davis and Hain¹⁷),

¹⁷ Urinary Antisepsis: The Antisoptic Properties of Normal Dog Urine, Jour. Urol., 1018, ii, 309.

which shows that normal dog and rabbit urine, for undetermined reasons, may occasionally act as an unfavorable culture medium for the colon bacillus and may even kill this organism after several hours. Each experiment was therefore accurately controlled by a specimen of urine obtained just before administration of the drug and inoculated and subjected to identically the same conditions as those specimens obtained at intervals after injection.

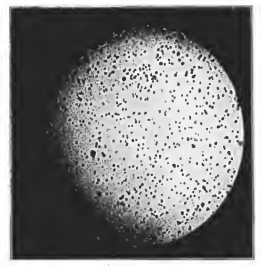


Fig. 6.—Photomicrograph (low power) of typical field in agar plate, showing a countless number of colonies and proving that the corresponding urine tube had contained an insufficient concentration of dye to prevent growth of the organism. The number of colonies in such a plate is designated in Table IV by the infinity sign ∞ .

Two samples (1 e.e. each) were transferred from each specimen of urine to sterile test-tubes, inoculated with B. coli and S. albus respectively (one loop of a twenty-four-hour broth enlture) and incubated twenty-four hours, after which 0.1 e.e. was transferred from each tube to melted agar and plated. This technic was identical with that described in a preceding paragraph for determining the antiseptic properties of the same dyes in voided human urine, with the exception that the dyes were not added to the urine but were excreted by the kidney. The plates were inspected after forty-eight hours. A plate showing no colonies (Fig. 4) or

very few colonies (Fig. 5) proved that particular specimen of urine to be antiseptic, while the presence of countless numbers of colonies (illustrated by Fig. 6 and designated in Table IV by the infinity sign ∞) proved that the organism had grown and developed and that the urine had acted as a favorable culture medium. This method, dependent upon observing the number of colonies in an agar plate, accurately determines whether the organism has developed or died during the incubation period and is not open to fallacy, as is the method dependent upon gross inspection of incubated urine.

Table IV summarizes the results of attempts to demonstrate antiseptic properties in rabbit urine following the intravenous administration of the 13 dyes shown by previous selection (Tables I. II and III) to be antiseptic, excreted and non-toxic in moderate dosage. The same table shows the results of similar experiments carried out with proflavine and acriflavine. As a previous publication has pointed out (Davis and White8), and as the above experiments verify, proflavin and acriflavin are experimentally successful in that antiseptic properties in the urine may be definitely demonstrated following intravenous administration. (The flavins giving these results were manufactured by the Boots Pure Drug Company, Nottingham, England.) Out of the entire list of 204 new dves. however, although preliminary experiment justified the final selection of thirteen, the properties of which indicated their possible value as internal urinary antiseptics, with none excepting proflavin and acriflavin, was it possible to cause the secretion of antiseptic urine by intravenous administration. In spite of the fact that these several selected dyes approached the ideal requirements (that is were antiseptic in urine, were excreted by the kidney and were relatively non-toxic), yet they failed at the final test when passed through the blood stream and kidney.

Conclusions.—1. There is no known drug ideally suited for the

purpose of internal urinary antisepsis.

2. Of a total of 204 anilin dyes investigated 61 were found to possess antiseptic properties in agar, and 28 of these were efficient

as antiseptics when added to voided urine.

3. As regards selective action against various organisms, this property was exhibited by no less than 44 dyes, in every case the colon bacillus proving more resistant than the staphylococci. There were only 24 which inhibited the colon bacillus in urine in a dilution of 1 to 1000.

4. There was almost no exception to the rule that antiseptic action was exhibited in higher dilution in alkaline urine than in acid urine. Attention is therefore called to the fact that these dyes are most efficient in urine of a reaction which renders urotropin inert.

5. The azo dyes give no promise of value, since of 37 of this group studied only 3 possessed antiseptic properties, and these only to a slight degree.

6. Of the triphenylmethanes many were antiseptic in high dilution in urine (some in dilution greater than 1 to 1,000,000). Of these, however, all but one were toxic and none was excreted by the This group is, nevertheless, worthy of further investigation.

7. Of 21 dyes of the xanthane group 3 were antiseptic in voided

urine, and 2 of these were excreted to a moderate degree.

8. Of 4 acridine dyes 2 were antiscritic in urine. Neither was excreted.

9. Of 9 dyes of the azine group 5 were antiscrtic in urine, and 3 of these (Safranin T, Safranin OW, Safranin MN) were excreted by the kidney with great rapidity and completeness and were non-toxic

in 20 mgm, per kilo dosage.

- 10. By a study of 204 anilin dyes, chosen at random, it has been possible to select 15 which are (a) antiseptic in urine, (b) excreted by the kidney and which are (c) relatively non-toxic. With only two of these, however (proflavin and acriflavin) was it possible to demonstrate the secretion of antiseptic urine following intravenous administration.8
- 11. Considering that rapid renal elimination of anilin dyes is not unusual; that there are not a few dyes, relatively non-toxic, which exert a bacteriostatic action when diluted to infinitesimal amounts in voided urine; and that out of 204 dyes it has been possible to select 15 which approach the ideal and 2 which are experimentally effective; it is within reasonable expectation that a dyc clinically suited for use as an internal urinary antiseptic may be discovered or synthesized. Experiments to date indicate that dyes of the triphenylmethane, xanthone, acridin and azin groups (particularly) the latter) give more promise of value.

THE FOURTH VENEREAL DISEASE, ULCERATIVE AND GAN-GRENOUS BALANOPOSTHITIS: WITH CASE REPORT.

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Judging by the dearth of case reports and the rare mention of the so-called "Fourth Venereal Disease," it would seem that the existence of this condition is not a matter of common knowledge among physicians. Since Corbus and Harris' first called it to the attention of the profession in this country in 1909 but two others have made case reports in the American journals.²³ Some of the

Jour. Am. Med. Assn., May 8, 1909, lii, 1474.

² Bond, S. P.: Urol. and Cut. Rev., 1919, xxiii, 211. ³ Ross, C. F.: Virginia Med. Monthly, xlv, 36.